

Thermal Transport across Graphite-Water Interfaces

D.E. Forero^{1,2}, Patrick Shamberger¹

¹ Thermal Sciences & Materials Branch, Materials and Manufacturing Directorate, Air Force Research Laboratory, Wright-Patterson AFB, OH 45433

² University of Dayton Research Institute, University of Dayton, Dayton, OH 45469

Motivation

- Need for thermal storage of large transient pulses of heat on USAF platforms
- Salt hydrates and graphitic foam composites offer high thermal energy storage capabilities and high thermal conductivities respectively
- Thermal resistance across graphite-hydrous salt interfaces is unknown

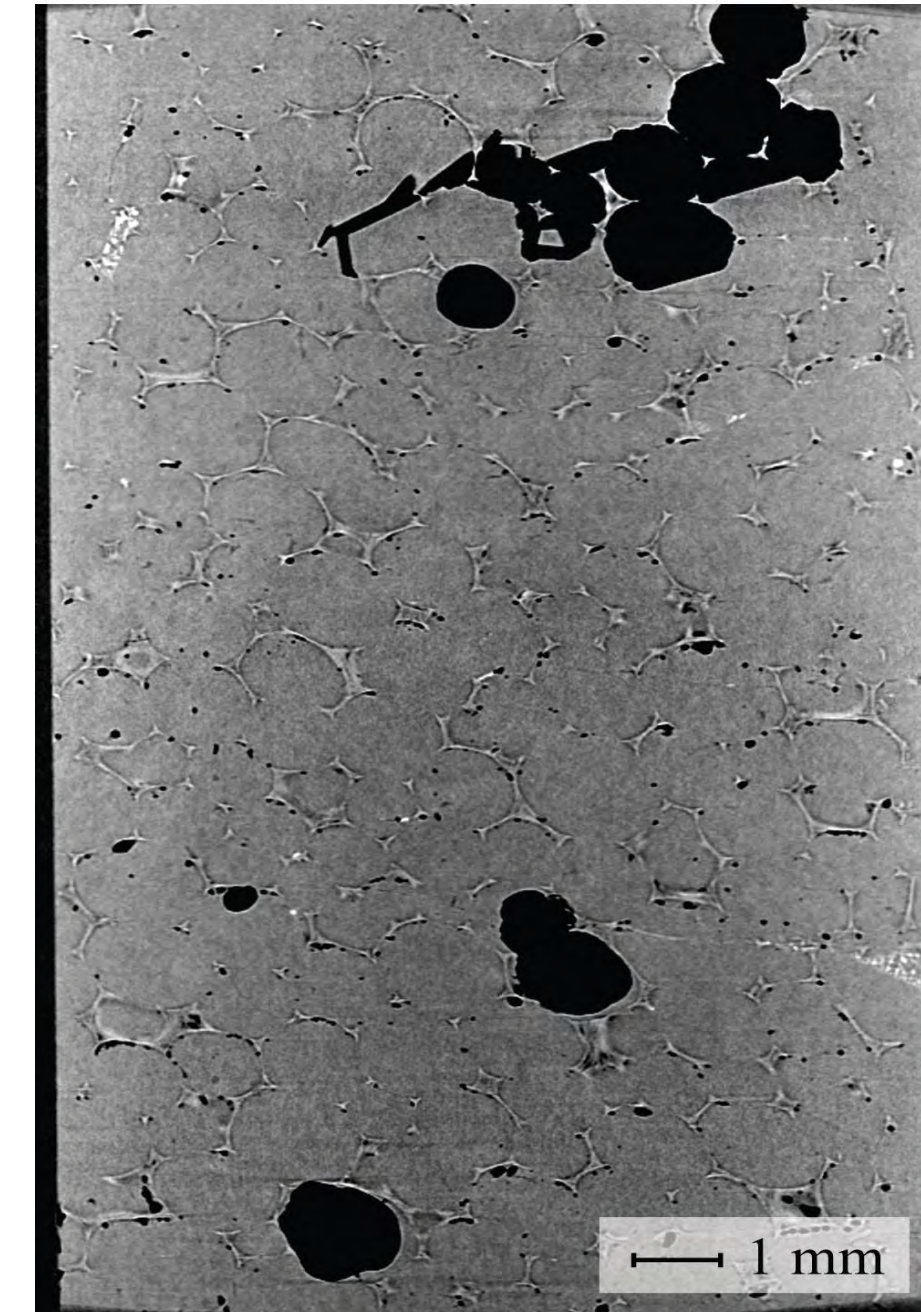


Fig.1 X-Ray Tomography Image of Graphitic foam wetting behavior

Objectives:

1. Analyze thermal transport across water graphite interfaces
2. Determine importance of thermal interfaces within composites

Graphite Wetting

- Graphite is naturally hydrophobic
- This is problematic when making a graphite/salt hydrate composite
- Solutions Considered
 - UV-Ozone treatment
 - Thin SiO_x layer
 - DOW Corning Q2-5211 Super wetting Agent (Commercial non-ionic silicone surfactant)
- This study focused on using Dow Super wetting (SW) agent due to its ease of use

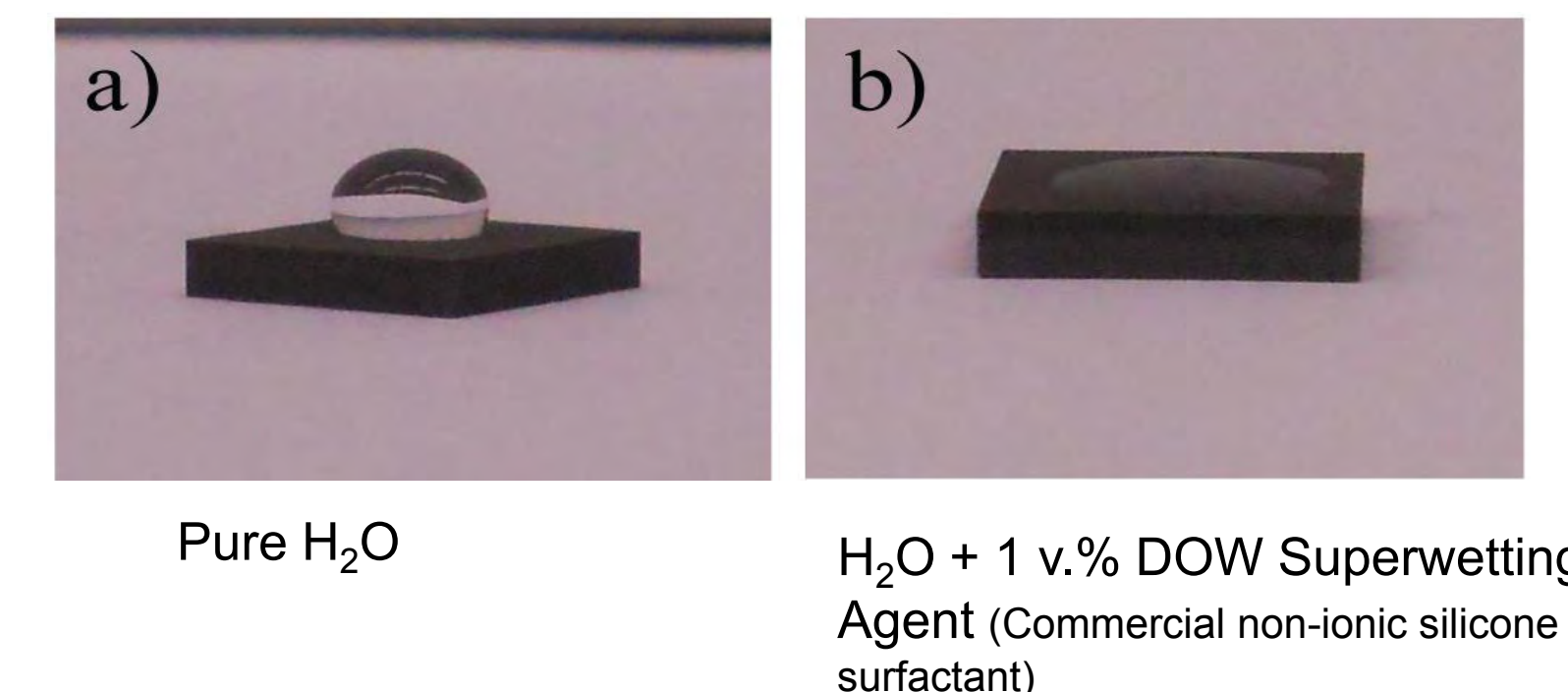


Fig.2 Superwetting agent effects on graphite wetting

Experimental Approach

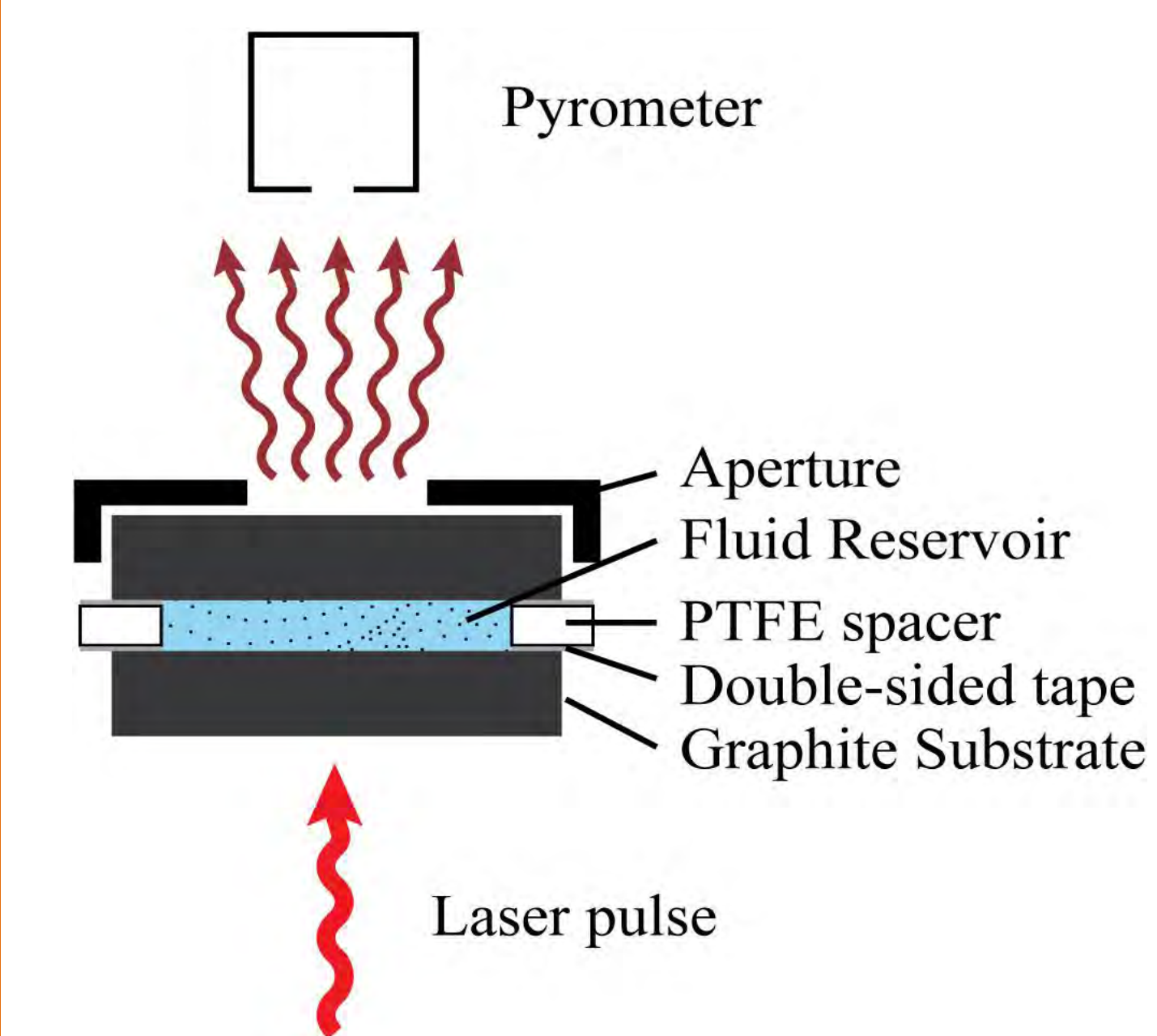


Fig. 3 Graphite-Water-Graphite stack set up.

- Thin water (36 μ l) layer sandwiched between two graphite substrates
- Water enclosed by Teflon/vacuum grease or Teflon/double-sided tape.
- Used Netzsch LFA 457 to conduct transient measurements
- Independently measured values were used for substrate properties
- Model fit resulted in correlation coefficients of .995 and higher.
- Data was best fit by using a model accounting for heat loss.

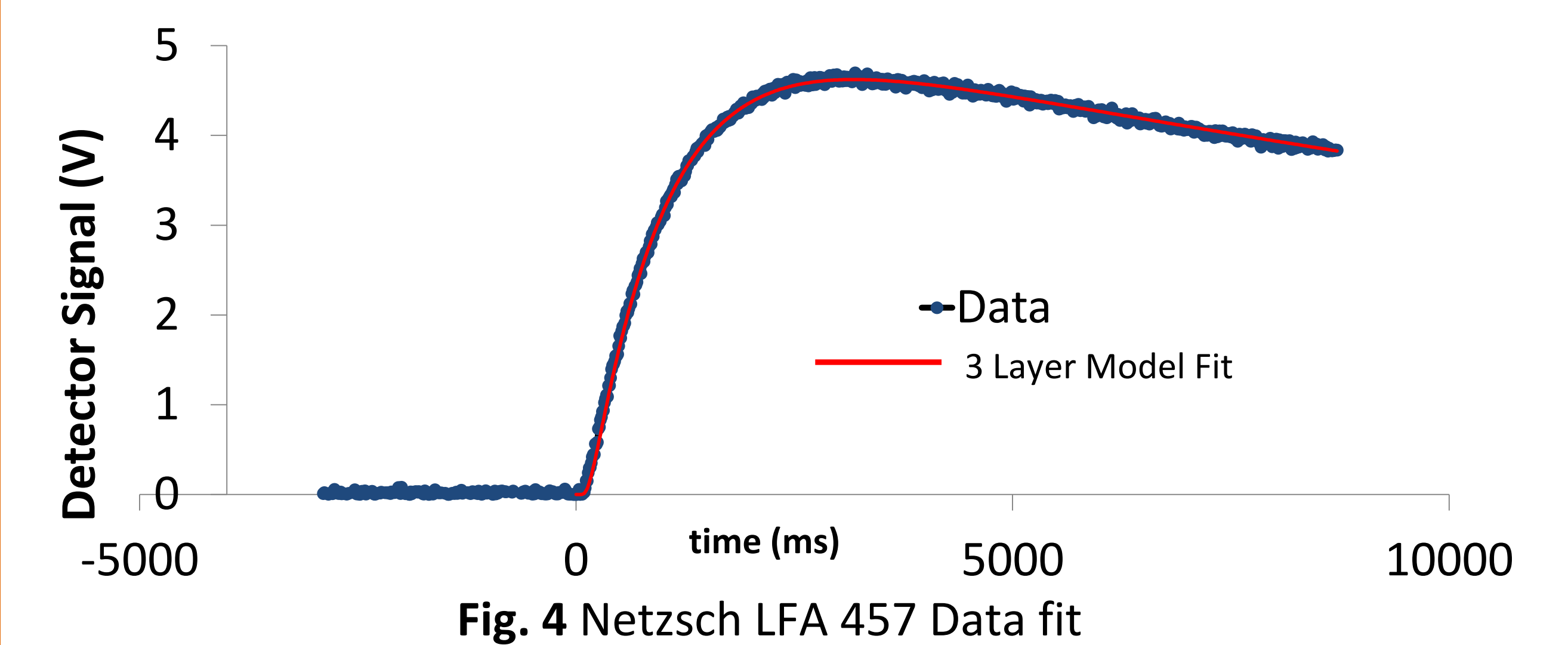


Fig. 4 Netzsch LFA 457 Data fit

Results

Effect of Thickness

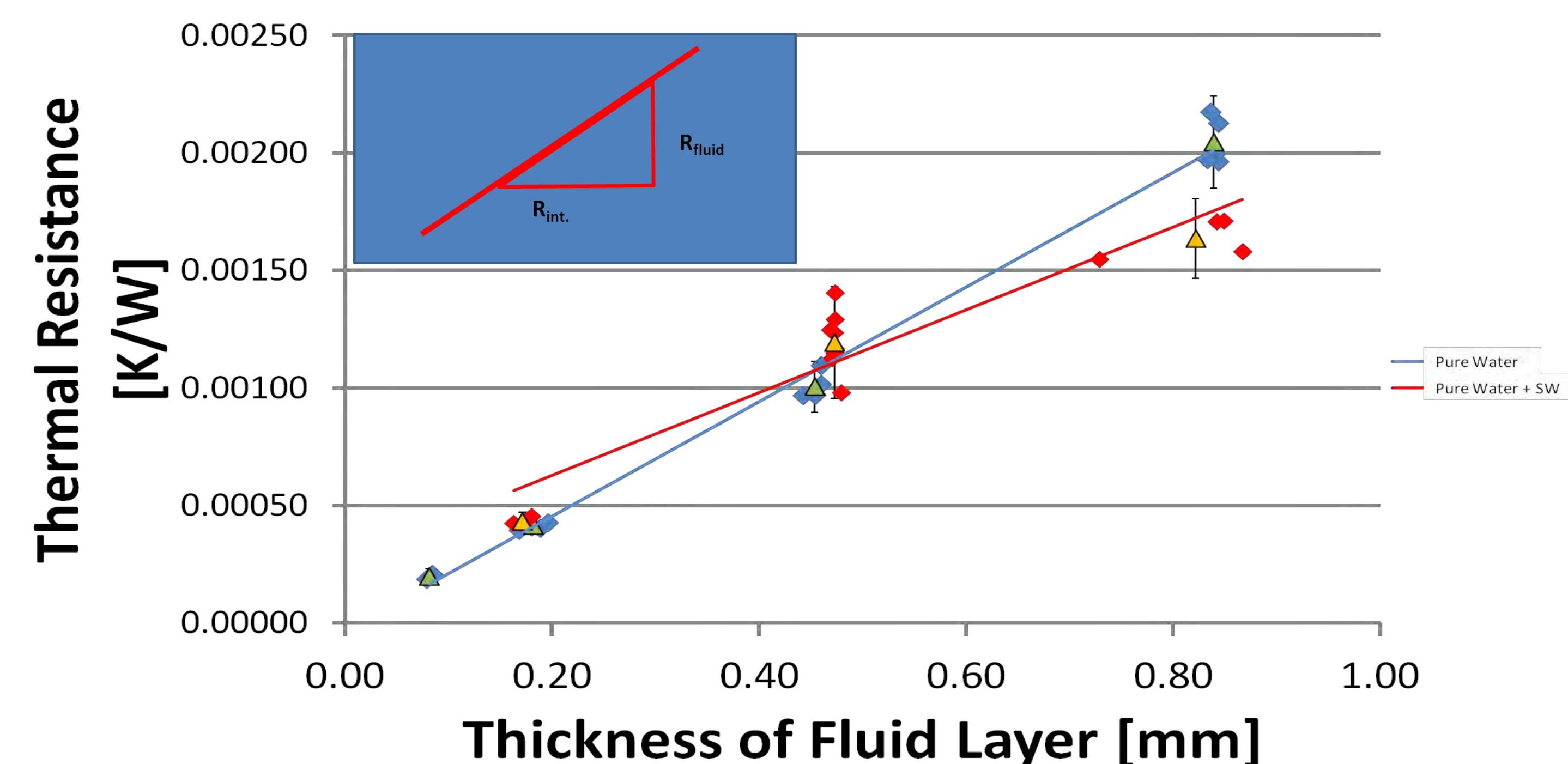


Fig.7 Thermal Resistance vs. Thickness of Fluid Layer

- $R_{\text{interface}} < 0.14 \text{ K}\cdot\text{cm}^2/\text{W}$ (at 95 % CI)
- $R_{\text{interface}} \ll R_{\text{fluid}} (\sim 10 \text{ K}\cdot\text{cm}^2/\text{W})$
- Previous measurements:¹
 $R_{\text{H}_2\text{O interface}} = 0.03 \text{ to } 0.2 \text{ K}\cdot\text{cm}^2/\text{W}$

¹ Ge, Z., Cahill, D.G., Brum, F.V., (2006), Thermal Conductance of Hydrophilic and Hydrophobic Interfaces, Phys. Rev. Lett. 96, 186101.

Effect of SW Concentration

- Concentration dependence on diffusivity was analyzed
- In all cases effective diffusivity values were lower than accepted literature values for bulk water
 - Difference from NIST value may be due to:
 1. Interpreting data with complex 3-Layer model
 2. Complexity of sample geometry
 3. Convection in fluid layer

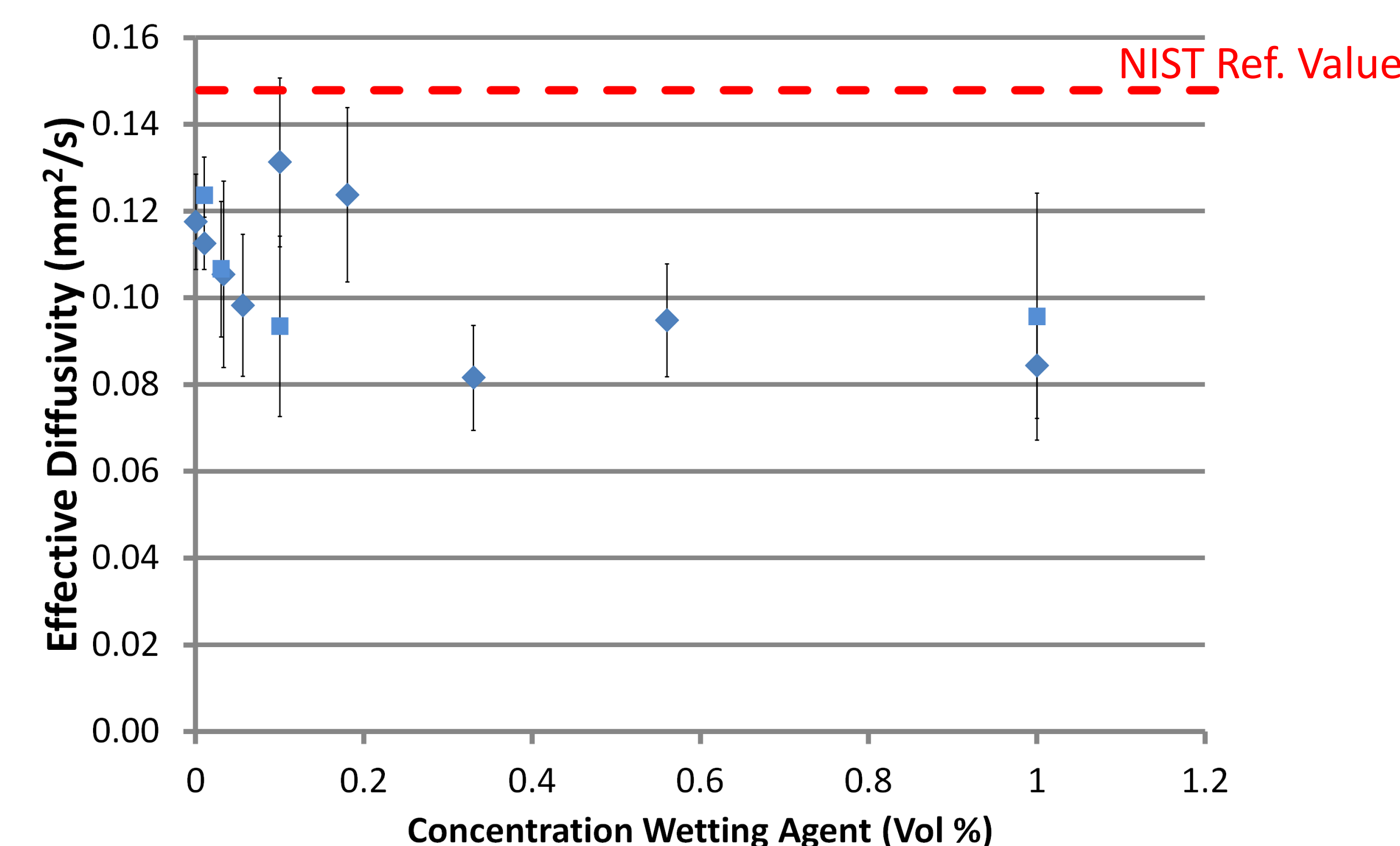


Fig.6 Effective Diffusivity vs. Concentration %

Conclusions

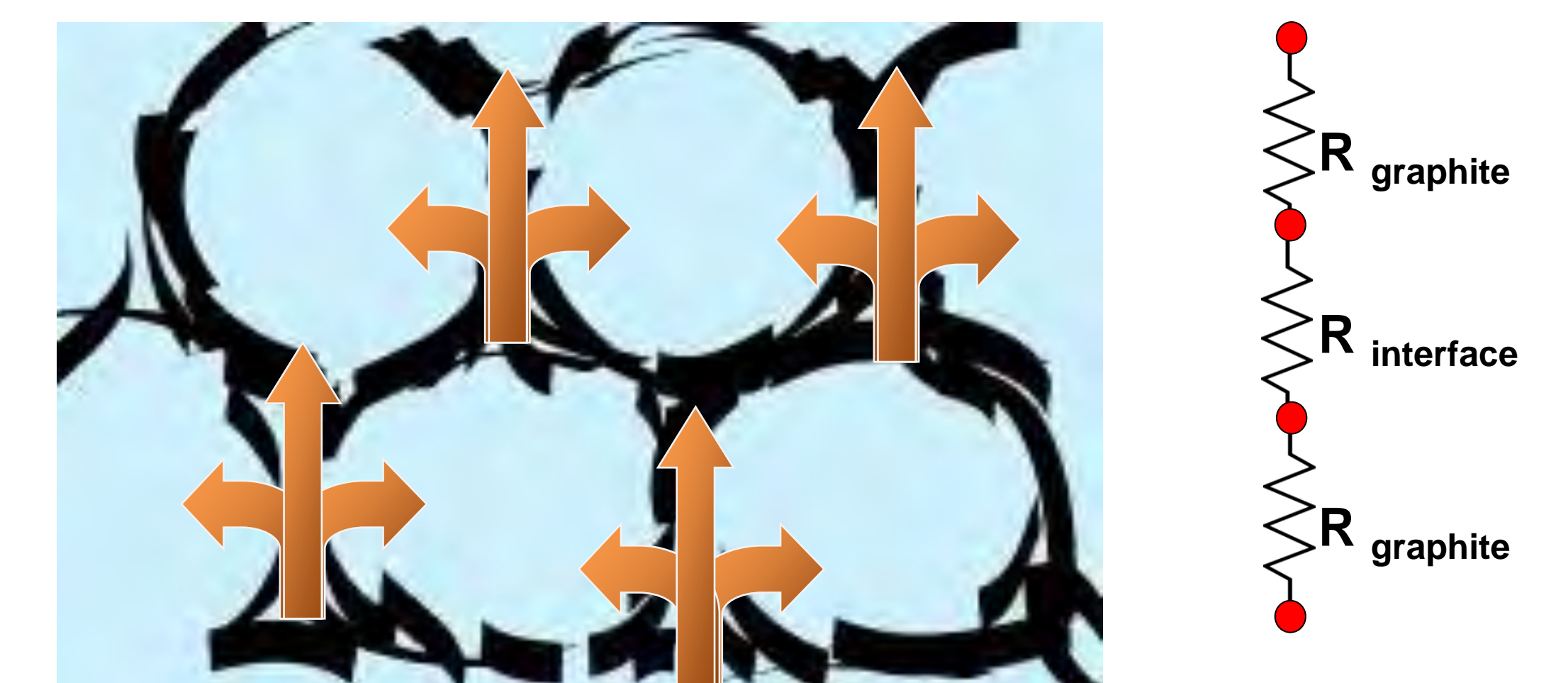


Fig.7 Heat transfer in graphitic foam

- In this case, $R_{\text{interface}} (\sim 0.10 \text{ K}\cdot\text{cm}^2/\text{W}) \ll R_{\text{fluid}} (\sim 10 \text{ K}\cdot\text{cm}^2/\text{W})$
- $R_{\text{interface}} (\text{Pure Water}) \sim R_{\text{interface}} (\text{Pure Water} + \text{SW})$
- Therefore interfaces are negligible when dealing with heat transfer within composites
- Use of surfactants proved successful in increasing wetting ability of graphite